

# Cambridge International AS & A Level

CANDIDATE NAME			
CENTRE NUMBER		CANDIDATE NUMBER	
CHEMISTRY			9701/42
Paper 4 A Leve	el Structured Questions		May/June 202
			2 hours
You must answ	ver on the question paper.		

#### **INSTRUCTIONS**

Answer all questions.

You will need: Data booklet

- Use a black or dark blue pen. You may use an HB pencil for any diagrams or graphs.
- Write your name, centre number and candidate number in the boxes at the top of the page.
- Write your answer to each question in the space provided.
- Do not use an erasable pen or correction fluid.
- Do **not** write on any bar codes.
- You may use a calculator.
- You should show all your working, use appropriate units and use an appropriate number of significant figures.

#### **INFORMATION**

- The total mark for this paper is 100.
- The number of marks for each question or part question is shown in brackets [ ].

This document has **24** pages. Any blank pages are indicated.

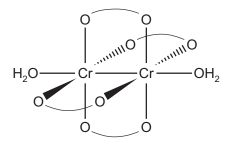
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## Answer **all** the questions in the spaces provided.

(a) An aqueous solution	of chromium( $ m III$ ) contains the gree	en $[Cr(H_2O)_6]^{3+}$ complex ion.
(i) Complete the ele	ectronic configuration of an isolated	d, gaseous Cr³+ ion.
1s²		[1
(ii) Define the term of	complex ion.	
		[1
2 102 1	vs some similar chemical properties $_6]^{3+}$ are reacted separately with e	s to $[Co(H_2O)_6]^{2+}(aq)$ . either NaOH(aq), $H_2O_2(aq)$ , or excess
Use this information	and the <i>Data Booklet</i> to suggest e of reaction taking place in each o	the formula of the chromium species case.
reagent added to [Cr(H <sub>2</sub> O) <sub>6</sub> ] <sup>3+</sup> (aq)	formula of chromium species formed	type of reaction
NaOH(aq)		
H <sub>2</sub> O <sub>2</sub> (aq)		
an excess of NH₃(aq)		
		[5]
colours.	O <sub>2</sub> CCH <sub>3</sub> ) <sub>4</sub> (H <sub>2</sub> O) <sub>2</sub> ] are both complexurs of these complexes are differer	lphaes of chromium( $ m II$ ) and have different
		[2 <sup>'</sup>

1

(d) The structure of  $[Cr_2(O_2CCH_3)_4(H_2O)_2]$  is shown. Ethanoate ions act as ligands in this complex. The ethanoate ligand,  $CH_3CO_2^{-}$ , is shown as  $O_2^{-}$ O.



	(i)	Water and ethanoate ions behave as different types of ligand in this complex.
		Suggest an explanation for this statement.
		[1
	(ii)	Deduce the coordination number of Cr and the geometry around each Cr atom in this structure.
		coordination number
		geometry around Cr atom[1
	(iii)	State the type of bond between the two atoms in the Cr–Cr bond.
		[1
(e)	The	$e [Cr_2(O_2CCH_3)_4(H_2O)_2]$ complex reacts with aqueous acid to form $Cr^{2+}(aq)$ ions.
	Cr <sup>2</sup>	$^{+}$ (aq) ions react with $O_2$ (aq) under acidic conditions. $Cr^{3+}$ (aq) ions are formed.
	Use	e the Data Booklet to answer the following questions.
	(i)	Construct an ionic equation for the reaction of Cr <sup>2+</sup> (aq) with O <sub>2</sub> (aq) under acidic conditions
		[2
	(ii)	Calculate $E_{coll}^{\bullet}$ for the reaction in (e)(i).

 $E_{\text{cell}}^{\bullet} = \dots V [1]$ 

[Total: 15]

(a)	Sta	te and explain the trend observed in the thermal stability of the Group 2 nitrates.
		[3]
(b)	(i)	Lead(II) nitrate, $Pb(NO_3)_2$ , decomposes on heating in a similar manner to the Group 2 nitrates.
		Write an equation for the decomposition of lead(II) nitrate.
		[1]
	(ii)	Suggest how the ease of decomposition of $Pb(NO_3)_2$ would compare to that of $Ba(NO_3)_2$ . Explain your answer. You may find it useful to refer to the <i>Data Booklet</i> .
		[1]
(c)	(i)	Barium ethanedioate, ${\rm BaC_2O_4}$ , decomposes on heating to produce barium oxide and a mixture of two different gases.
		Construct an equation for the decomposition of barium ethanedioate.
		[1]

2

(ii) An impure sample of  $BaC_2O_4$ , of mass 0.500 g, is added to 50.0 cm<sup>3</sup> of 0.0200 mol dm<sup>-3</sup> acidified  $MnO_4^-$ (aq), an excess. A redox reaction takes place and all the  $BaC_2O_4$  reacts.

The resulting solution, containing unreacted acidified  $MnO_4^-$ , is titrated with 0.0500 mol dm<sup>-3</sup>  $Fe^{2+}(aq)$ .

The end-point is reached when 30.40 cm<sup>3</sup> of 0.0500 mol dm<sup>-3</sup> Fe<sup>2+</sup>(aq) has been added.

$$C_2O_4^{2-} \rightleftharpoons 2CO_2 + 2e^ MnO_4^- + 8H^+ + 5e^- \rightleftharpoons Mn^{2+} + 4H_2O$$
 $Fe^{2+} \rightleftharpoons Fe^{3+} + e^-$ 

Calculate the percentage by mass of  $BaC_2O_4$  in the 0.500 g impure sample. Show your working.

 $[M_r: BaC_2O_4, 225.3]$ 

percentage by mass of 
$$BaC_2O_4 = \dots$$
 [4]

(d) Barium hydroxide,  $Ba(OH)_2$ , is completely dissociated in aqueous solution.

Calculate the pH of 0.120 mol dm<sup>-3</sup> Ba(OH)<sub>2</sub>(aq) at 298 K.

[Total: 12]

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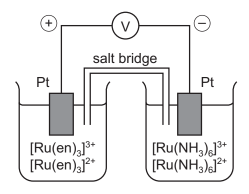
3	(a)	(i)	Define	the term	standard	electrode	potential
•	14/	111		tile terrii	otarraara	CICOLI CGC	poteritiai

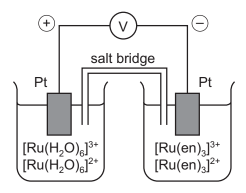
[2]	

Three redox systems,  $\bf A$ ,  $\bf B$  and  $\bf C$ , are shown. The ligand 1,2-diaminoethane,  $H_2NCH_2CH_2NH_2$ , is represented by en.

Α	$[Ru(H_2O)_6]^{3+} + e^- \rightleftharpoons [Ru(H_2O)_6]^{2+}$
В	$[Ru(NH_3)_6]^{3+} + e^- \rightleftharpoons [Ru(NH_3)_6]^{2+}$
С	$[Ru(en)_3]^{3+} + e^- \rightleftharpoons [Ru(en)_3]^{2+}$

Two electrochemical cells are set up to compare the standard electrode potentials,  $E^{\circ}$ , of three half-cells. The diagrams show the relative potential of each electrode.





(ii) Use this information to complete the table by adding the labels A, B and C to deduce the order of  $E^{\circ}$  for the three half-cells.

E <sup>e</sup>	redox system
most negative	
<b>1</b>	
least negative	

[1]



(iii) The complex  $[Ru(en)_3]^{3+}$  shows stereoisomerism. The ligand en is bidentate.

		i		:		
		isomer 1		isomer 2		
		 Ru		 Ru		
	type of stered	oisomerism				[;
)	conditions. Use the <i>Data</i>		late the $E_{\mathrm{cell}}^{\bullet}$ . Dec	alf-cell and a Ag <sup>+</sup> /A duce the direction -cells.		
)	conditions.  Use the <i>Data</i> through the v	Booklet to calcu	late the $E_{\mathrm{cell}}^{\bullet}$ . Dec	duce the direction		
)	conditions.  Use the <i>Data</i> through the v $E_{\text{cell}}^{\bullet} = \dots$	Booklet to calcuroltmeter betwee	late the $E_{\mathrm{cell}}^{ullet}$ . Deon these two half	duce the direction	of electron fl	ow in the wi
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$	Booklet to calcuroltmeter betwee	late the $E_{\mathrm{cell}}^{\bullet}$ . Dec n these two half	duce the direction -cells.	of electron fl	ow in the wi
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$ direction of example $E_{\text{cell}}^{\text{e}} = E_{\text{cell}}^{\text{e}} $	Booklet to calcuroltmeter between V  lectron flow from the dot to the Ag*/Ag	late the $E_{\text{cell}}^{\bullet}$ . Deen these two half-	duce the direction -cells.	of electron fl	ow in the win
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$ direction of example $E_{\text{cell}}^{\text{e}} = E_{\text{cell}}^{\text{e}} $	Booklet to calcuroltmeter between V  lectron flow from the dot to the Ag*/Ag	late the $E_{\text{cell}}^{\bullet}$ . Deen these two half-	duce the direction -cells. to	of electron fl	ow in the win
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$ direction of example $E_{\text{cell}}^{\text{e}} = E_{\text{cell}}^{\text{e}} $	Booklet to calcuroltmeter between V  lectron flow from the dot to the Ag*/Ag effect of this add	late the $E_{\rm cell}^{\bullet}$ . Deen these two half-half-cell in <b>(b)(i)</b> ition on the $E_{\rm cell}$ .	duce the direction cells.  to  Place a tick (✓) ir	of electron fl	ow in the wi
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$ direction of example $E_{\text{cell}}^{\text{e}} = E_{\text{cell}}^{\text{e}} $	Booklet to calcuroltmeter between V lectron flow from led to the Ag*/Ag leffect of this add less positive	late the $E_{\rm cell}^{\bullet}$ . Deen these two half-half-cell in <b>(b)(i)</b> ition on the $E_{\rm cell}$ .	duce the direction cells.  to  Place a tick (✓) ir	of electron fl	ow in the win
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$ direction of example $E_{\text{cell}}^{\text{e}} = E_{\text{cell}}^{\text{e}} $	Booklet to calcuroltmeter between V lectron flow from led to the Ag*/Ag leffect of this add less positive	late the $E_{\rm cell}^{\bullet}$ . Deen these two half-half-cell in <b>(b)(i)</b> ition on the $E_{\rm cell}$ .	duce the direction cells.  to  Place a tick (✓) ir	of electron fl	ow in the win
	conditions.  Use the <i>Data</i> through the volume $E_{\text{cell}}^{\text{e}} = \dots$ direction of example $E_{\text{cell}}^{\text{e}} = E_{\text{cell}}^{\text{e}} $	Booklet to calcuroltmeter between V lectron flow from led to the Ag*/Ag leffect of this add less positive	late the $E_{\rm cell}^{\bullet}$ . Deen these two half-half-cell in <b>(b)(i)</b> ition on the $E_{\rm cell}$ .	duce the direction cells.  to  Place a tick (✓) ir	of electron fl	ow in the wi

[Turn over

(c)		Silver bromide, AgBr, dissolves in an aqueous solution of $S_2O_3^{2-}$ ions to form the complex ion $[Ag(S_2O_3)_2]^{3-}$ . The $S_2O_3^{2-}$ ions act as monodentate ligands.							
	equ	ıilibrium	1 A	gBr(s) + 2S <sub>2</sub> O <sub>3</sub> <sup>2-</sup> (aq	$\Rightarrow$ [Ag(S <sub>2</sub> O <sub>3</sub>	) <sub>2</sub> ] <sup>3-</sup> (aq) + B	r⁻(aq)		
	(i)	Define	the ter	m <i>ligand</i> .					
							[1]		
	(ii)	Write a	an expre	ession for the equilib	rium constant,	$K_{c}$ , for equilib	rium 1.		
		<b>K</b> <sub>c</sub> =					[4]		
(	(iii)	Some	additior	nal data are given ab	out the dissolut	tion of AgBr ir	[1] n S <sub>2</sub> O <sub>3</sub> ²-(aq).		
				equilibrium cons	tant	numerical v	/alue		
			S	solubility product, $K_{sp}$	, of AgBr	5.4 × 10	-13		
			stabi	lity constant, $K_{\text{stab}}$ , of	[Ag(S <sub>2</sub> O <sub>3</sub> ) <sub>2</sub> ] <sup>3-</sup>	2.9 × 10	13		
(d)	The give		ical val	lues for the stability			nits[2] er silver(I) complexes are		
				silver(I) complex	numerical va	alue of K <sub>stab</sub>			
				$[Ag(CN)_2]^-$	5.3 ×	10 <sup>18</sup>			
				$[Ag(NH_3)_2]^+$	1.6 ×	10 <sup>7</sup>			
	CN <sup>-</sup>	In aqueous solution containing $Ag^+$ is added to a solution containing equal conc $CN^-(aq)$ , $NH_3(aq)$ and $S_2O_3{}^{2-}(aq)$ . The mixture is left to reach equilibrium.							
		ulting m	ixture. I	Explain your answer.			$[Ag(S_2O_3)_2]^{3-}$ present in the		
				>			ncentration		
							[2]		
							[Total: 15]		

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(a) (i)	Define the term <i>lattice energy</i> .
	[2]
(ii)	Use the following data to calculate a value for the enthalpy change of solution of copper(II) chloride, ${\rm CuC}l_2(s)$ . You might find it helpful to construct an energy cycle.
	enthalpy change of hydration of $Cl^- = -378  \text{kJ}  \text{mol}^{-1}$ enthalpy change of hydration of $Cu^{2+} = -2099  \text{kJ}  \text{mol}^{-1}$ lattice energy of $CuCl_2(s) = -2824  \text{kJ}  \text{mol}^{-1}$
	enthalpy change of solution of $CuCl_2(s) = \dots kJ mol^{-1}$ [2]
(iii)	The enthalpy change of hydration of Ca <sup>2+</sup> is –1579 kJ mol <sup>-1</sup> .
	Use the <i>Data Booklet</i> to suggest why there is a big difference in the values of $\Delta H_{\rm hyd}$ for Ca <sup>2+</sup> and Cu <sup>2+</sup> .
	[2]
(b) (i)	Identify the substances formed at the anode and at the cathode during the electrolysis of saturated $CaCl_2(aq)$ .
	at the anode
	at the cathode[1]
(ii)	Calcium can be produced by the electrolysis of molten calcium chloride, ${\rm CaC}l_2({\rm I}).$
	Calculate the mass, in g, of Ca formed when a current of 0.75A passes through $CaCl_2(I)$ for 60 minutes. [ $A_r$ : Ca, 40.1]
	mass of Ca = g [2]

(c) (i)		Explain what is meant by the term <i>entropy of a system</i> .	
			[1]

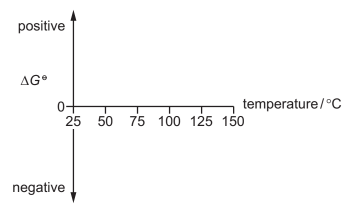
(ii) Place one tick ( $\checkmark$ ) in each row of the table to show the sign of each entropy change,  $\triangle S$ .

process	$\Delta S$ is negative	$\Delta S$ is zero	$\Delta S$ is positive
NaC <i>l</i> dissolving in water			
water solidifying to ice			

[1]

(iii) The evaporation of one mole of water has a standard Gibbs free energy change,  $\Delta G^{\circ}$ , of +8.6 kJ at 25 °C.

Sketch a graph on the axes to show how  $\Delta G^{\circ}$  changes for this process between 25 °C and 150 °C at 101 kPa.



[2]

(d) The reaction between A and B is feasible at low temperatures but is **not** feasible at high temperatures.

$$A + B \rightleftharpoons C + D$$

Deduce the signs of  $\Delta H$  and  $\Delta S$  for this reaction and explain why the feasibility changes with temperature.

sign of $\Delta H = \dots$	sign of $\Delta S$ =

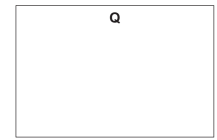
.....[2]

[Total: 15]

5	(a)	Describe and explain the relative basicities	of phenylamine	, ethylamine and	4-nitrophenylamine.

	>	>	
most basic		least basic	

(b) The dye  ${\bf R}$  can be synthesised from 4-nitrophenylamine in two steps.

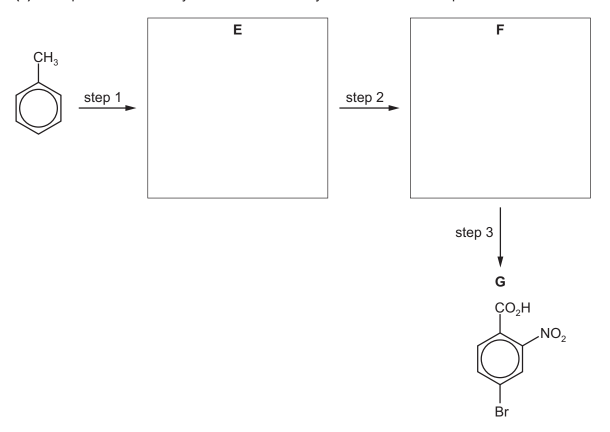


$$R$$
 $O_2N$ 
 $N$ 
 $N$ 
 $N$ 
 $N$ 
 $N$ 

- (i) Deduce and draw the structure of the organic salt **Q** in the box. [1]
- (ii) Suggest reagents and conditions for step 1 and 2 in (b).

step 1	
step 2	
•	[2]

(c) Compound G can be synthesised from methylbenzene in three steps.



(i)	Give the systematic name of compound <b>G</b> .
` '	- ,

·	4.7
l'annual de la companya de la compa	1

- (ii) Deduce the identities of **E** and **F** and draw their structures in the boxes. [2]
- (iii) Suggest reagents and conditions for each of steps 1 to 3 in (c).

step 1	 
step 2	
step 3	 
	[3]

[Total: 13]

6 (a) There are four possible structural isomers of  $C_8H_{10}$  that contain a benzene ring.

Draw the **skeletal** formulae of the four structural isomers in the appropriate boxes. The number of peaks observed in the carbon-13 (<sup>13</sup>C) NMR spectrum of each compound is given.

three peaks in <sup>13</sup>C NMR

four peaks in <sup>13</sup>C NMR

isomer 3

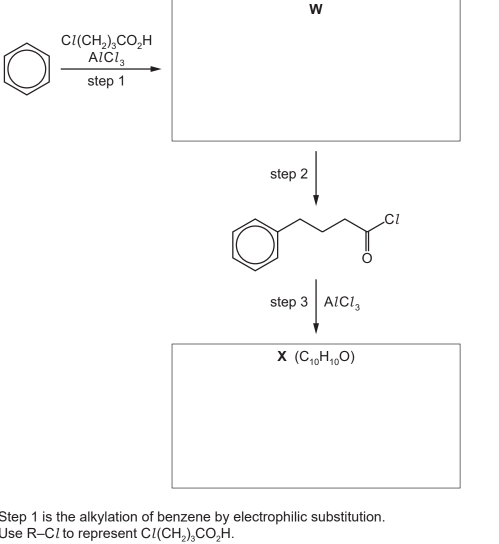
isomer 4

five peaks in <sup>13</sup>C NMR

six peaks in <sup>13</sup>C NMR

[4]

**(b)** A three-step synthesis of  $\mathbf{X}$  ( $C_{10}H_{10}O$ ) from benzene is suggested as shown.



(i)	Step 1 is the alkylation of benzene by electrophilic substitution.
	Use R–C $l$ to represent $Cl(CH_2)_3CO_2H$ .

Write an equation for the formation of an electrophile from R-Cl and  $AlCl_3$ .

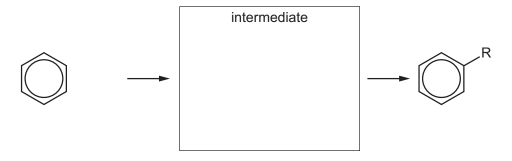
- Deduce and draw the structures of **W** and **X** in the boxes. [2]
- (iii) Suggest the reagents and conditions for step 2.



(iv) Complete the mechanism for the reaction of benzene with the electrophile formed in (b)(i).

Include all relevant charges and curly arrows showing the movement of electron pairs.

Draw the structure of the intermediate.



[3]

[Total: 11]

7 (a) In aqueous solution, chlorine dioxide,  $ClO_2$ , reacts with hydroxide ions as shown.

$$2\text{C}l\text{O}_2 \text{ + 2OH}^- \rightarrow \text{C}l\text{O}_3^- \text{ + C}l\text{O}_2^- \text{ + H}_2\text{O}$$

A series of experiments is carried out using different concentrations of  $\mathrm{C}l\mathrm{O}_2$  and  $\mathrm{OH}^-$ . The table shows the results obtained.

experiment	$[ClO_2]$ /moldm $^{-3}$	[OH <sup>-</sup> ] /moldm <sup>-3</sup>	initial rate /moldm <sup>-3</sup> min <sup>-1</sup>
1	0.020	0.030	7.20 × 10 <sup>-4</sup>
2	0.020	0.120	$2.88 \times 10^{-3}$
3	0.050	0.030	4.50 × 10 <sup>-3</sup>

(i)	Explain the term <i>order of reaction</i> .
	[1
(ii)	Use the data in the table to determine the order of reaction with respect to each reactant $\rm C\it lO_{2}$ and $\rm OH^{-}$ .
	Explain your reasoning.
	[2
(iii)	Use your answer to (a)(ii) to construct the rate equation for this reaction.
	rate =[1
(iv)	Use your rate equation and the data from experiment 1 to calculate the rate constant, $k$ for this reaction. Include the units of $k$ .

k =	units	[2]
$\kappa = \dots$	uiilo	[-]

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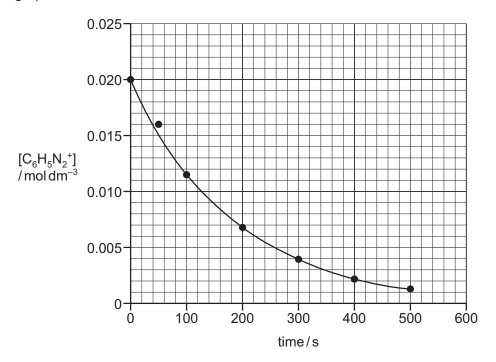


Question 7 continues on the next page.



**(b)** The decomposition of benzenediazonium ions,  $C_6H_5N_2^+$ , using a large excess of water, is a first-order reaction.

The graph shows the results obtained.



(i) Draw the structure of the organic product formed in this reaction.

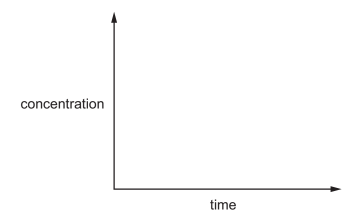
[1]

(ii) Use the graph to determine the rate of reaction at 100 s. Show your working.

rate = .....  $mol dm^{-3} s^{-1}$  [1]

(c) Sketch a concentration—time graph for a **zero-order** reaction.

Use your graph to suggest how successive half-lives for a zero-order reaction vary as the concentration of a reactant decreases. Indicate this by placing a tick ( $\checkmark$ ) in the appropriate box in the table.



successive half-lives decrease	no change in successive half-lives	successive half-lives increase

[1]

[Total: 9]

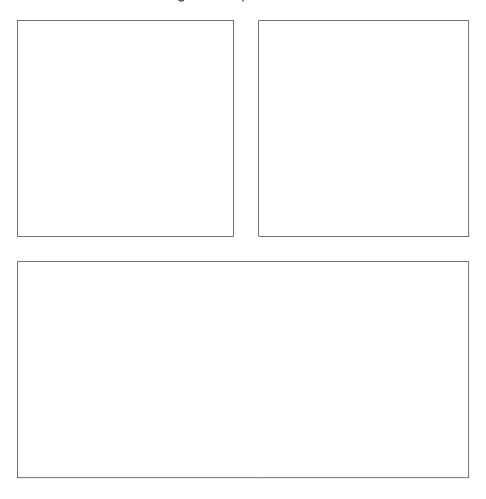
8	(a)	State and explain the relative rate of aryl chlorides.	hydrolysis of acyl chlorides, alkyl chlorides and
		fastest	>slowest

## (b) The drug remifentanil is shown.

remifentanil

Remifentanil is **completely** hydrolysed under acidic conditions. Three different organic compounds are formed.

Draw the structures for these organic compounds in the boxes.

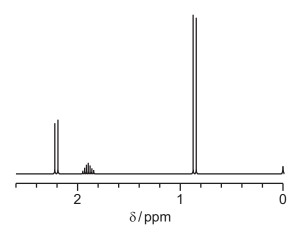


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(c) Compound Y,  $C_5H_{10}O_2$ , reacts with  $Na_2CO_3(aq)$  to evolve bubbles of gas. The proton (<sup>1</sup>H) NMR spectrum of compound Y in  $D_2O$  is shown.



(i) Use this information to suggest a structure for Y.

[1]

(ii) Use the *Data Booklet*, the proton (¹H) NMR spectrum and your answer to (c)(i) to complete the table.

chemical shift (δ)	environment of proton	splitting pattern	number of <sup>1</sup> H atoms responsible for the peak
0.95			
1.90			
2.20			

[3]

[Total: 10]



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